

Unboxing mutations: Connecting mutation types with evolutionary consequences

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Abstract

A key step in understanding the genetic basis of different evolutionary outcomes (e.g., adaptation) is to determine the roles played by different mutation types (e.g., SNPs, translocations and inversions). To do this we must simultaneously consider different mutation types in an evolutionary framework. Here, we propose a research framework that directly utilizes the most important characteristics of mutations, their population genetic effects, to determine their relative evolutionary significance in a given scenario. We review known population genetic effects of different mutation types and show how these may be connected to different evolutionary outcomes. We provide examples of how to implement this framework and pinpoint areas where more data, theory and synthesis are needed. Linking experimental and theoretical approaches to examine different mutation types simultaneously is a critical step towards understanding their evolutionary significance.

KEYWORDS

adaptation, distribution of fitness effects, mutation, mutation rate, population genetics, recombination, speciation, structural variant

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1 | INTRODUCTION

Understanding the genetic underpinnings of processes such as adaptation and speciation is a major goal in evolutionary biology. This requires quantifying the number of involved loci and their distribution in the genome (e.g., genomic architecture) as well as the type of mutation and their associated statistical properties. There has been a recent focus on types of mutations (e.g., SNPs, translocations, inversions) and detailed discussions of the evolutionary significance of different mutation types abound in the literature (e.g., Choi & Lee, 2020; Faria et al., 2019; Katju & Bergthorsson, 2013). However, most of these reviews, as well as most theoretical and empirical studies, do not examine the full range of mutation types. In order to understand the relative evolutionary significance of different mutation types we must consider them simultaneously in an integrated framework.

From an evolutionary viewpoint, the most important characteristics of a mutation are its occurrence rate, its population genetic effects and how these may influence downstream evolutionary outcomes. Here, we propose a research framework that takes advantage of decades of population genetic research (Charlesworth & Charlesworth, 2010; Futuyma, 1986) to directly utilize the population genetic effects of different mutation types to determine their relative evolutionary significance (Figure 1a). We consider population genetic effects to be the impact of a mutation on the population genetic parameters: recombination rate, effective population size, selection and dominance coefficients (see Figure 1c for an overview of the population genetic effects of each mutation type). Although other characteristics, such as whether a mutation affects regulatory or coding regions, are also important for evolution (Hoekstra & Coyne, 2007), we concentrate here on characteristics that can be directly tied to mutation type.

Our framework combines what we term the “forward” and the “reverse” approach. Starting from mutation, the forward approach characterizes how often different types of mutations occur (Figure 1b) and their population genetic effects (Figure 1c), in order to predict their role in different evolutionary outcomes. The reverse approach starts with the evolutionary outcome and, using empirical data, determines the relative contributions of multiple mutation types in a systematic fashion. There is already a wealth of knowledge on the population genetic effects of different mutations (i.e., the forward approach; Dobigny et al., 2017; Elena et al., 1998; Korunes & Noor, 2019; Malik, 2009). In contrast, few studies examine multiple types of mutations simultaneously using the reverse approach. Thus, here we concentrate on the forward approach.

To outline the forward approach, we give a nonexhaustive summary of what is known about mutation rates and the population genetic effects of the different mutations, taking advantage of the extensive existing body of work (Choi & Lee, 2020; Faria et al., 2019; Katju & Bergthorsson, 2013). We stress that quantifying these effects, evaluating differences between mutation types, and determining the distribution of effect sizes will be a critical step forward

and suggest different ways that this can be accomplished. We discuss areas where more empirical data, theory and synthesis are needed for the forward approach, and highlight the importance of applying the reverse approach.

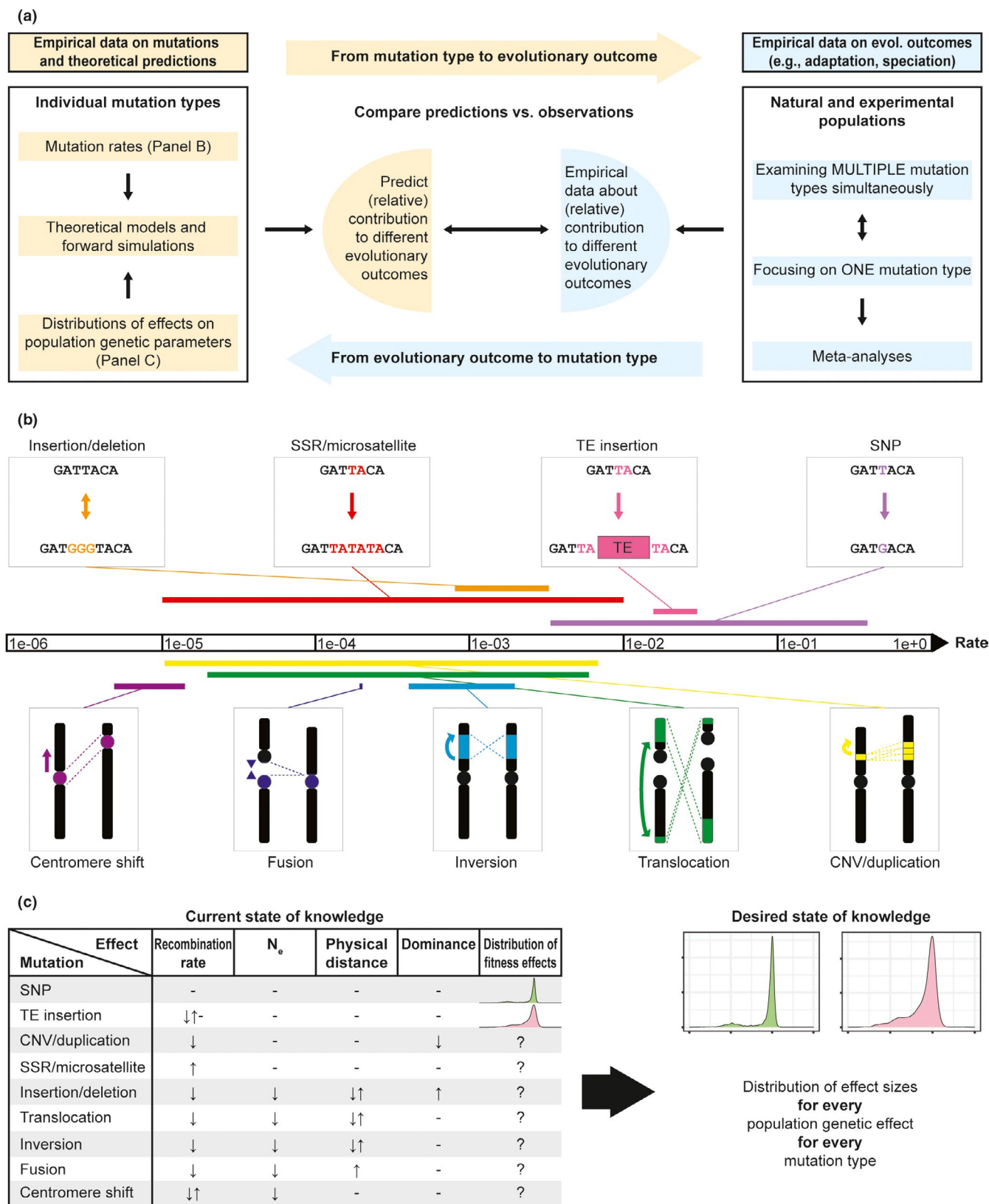
1.1 | Mutation rate

Mutation rate is a critical parameter when investigating the evolutionary impact of different types of mutations. It can be measured directly, by comparing the number of mutations in gametes, zygotes or offspring (Fu & Huai, 2003). For SNPs it can also be measured indirectly by comparing synonymous (usually presumed neutral) polymorphism data within and between closely related species. Estimates of the mutation rate vary greatly across taxa and mutation types (Figure 1b). However, different mutations vary with respect to their detectability (for example, SNPs are much more likely to be detected with short-read sequencing data than larger insertions or deletions [Ho et al., 2020]), which can lead to an underestimation of the mutation rate of different mutations. New methodologies can help ameliorate this issue. Specifically, long read data in combination with other methodologies such as linked read sequencing (Seo et al., 2016) and chromatin capture and sequencing techniques such as Hi-C (Wang et al., 2020) can facilitate the identification of large structural variants (Christmas et al., 2019). Thus, two key steps identified by our framework are to (i) further develop new methods that allow simultaneous detection of the different mutation types, and (ii) increase both the number as well as the taxonomic breadth of studies that directly estimate mutation rates and analyse multiple different mutation types within the same taxon.

2 | POPULATION GENETIC EFFECTS OF MUTATIONS: WHAT DO WE KNOW?

In the following sections, we discuss the effects of different mutation types on population genetic parameters such as recombination rate and effective population size. Note that we do not include chromosomal fissions due to lack of information, nor whole genome duplications as they have been covered extensively in previous reviews (e.g., Fox et al., 2020). While there is a wealth of theory that describes the evolutionary impact of mutations modifying population genetic parameters (Charlesworth & Charlesworth, 2017; Guerrero & Kirkpatrick, 2014; Kirkpatrick, 2010; Llaurens et al., 2015; Otto & Barton, 1997) these effects are rarely tied to a specific type of mutation and the entirety of effects of a single mutation are seldom taken into account. Mutations can also affect DNA methylation or chromatin state; however, as these are not standard population genetic parameters they are discussed separately in Box 1.

To make predictions about evolutionary outcomes, it is important to not just estimate the direction, but also the effect size of the changes in population genetic parameters (see Figure 1c: current vs.



desired state of knowledge). This can be done in two different ways: (i) By directly measuring the effect (e.g., recombination rate) empirically, or (ii) by incorporating information from molecular mechanisms

and empirical estimates, and feeding these into theoretical models to quantify these effects. While we mostly focus on (i) in the main text, we provide an example of (ii) (for recombination rate) in Box 2.

FIGURE 1 (a) Overview schematic illustrating forward and reverse approaches to determine the relative evolutionary significance of different mutation types. Forward approaches (yellow, from left to right) start from mutation rates and population genetic effects of mutations and feed these into theoretical models and simulations to generate predictions. Reverse approaches (blue, from right to left) start with evolutionary outcomes (e.g., adaptation) and examine the roles of different mutations simultaneously or single mutation types at a time, using data obtained either from natural populations or experimental evolution. The results from single mutation studies can be combined in a meta-analysis. The results from the reverse approach can then be compared with the predictions from the forward approach. (b) Overview of mutation types and their mutation rates. Mutation rates are per genome per generation, come from a wide range of taxa, and are taken from Beckmann et al. (2007), Brumfield et al. (2003), Ducos et al. (2007), Farlow et al. (2015), Feusier et al. (2019), Fu et al. (2010), Gemayel et al. (2010), Goerner-Potvin and Bourque (2018), Jarne and Lagoda (1996), Katju and Bergthorsson (2013), Maeda et al. (1991), Marriage et al. (2009), Marshall et al. (2008), Ossowski et al. (2010), Ramu et al. (2013), Rocchi et al. (2012), Schrider et al. (2013), Sung et al. (2016), Thuillet et al. (2002), Vendrell-Mir et al. (2019), Weng et al. (2019) and Yamaguchi and Mukai (1974). Please note that this is not an exhaustive overview and that actual ranges are probably larger. (c) Current and desired state of knowledge regarding the effects of mutation types on population genetic parameters. Current state of knowledge: Arrows indicate an increase (up) or decrease (down) while a dash indicates no known effect. For more details of the effects please refer to the *Population Genetic Effects of Mutations: What do we know?* section 2. Indels and inversions are assumed to be large enough to affect pairing at meiosis while smaller indels and inversions are expected to behave similarly to SNPs. The DFEs are schematics simulated from different statistical distributions and do not reflect real data. Desired state of knowledge: Colour and shape of schematic distributions (simulated) do not reflect real data or predictions

BOX 1 Beyond DNA sequence alterations

Many mutations affect not only the DNA sequence but also the local state of DNA methylation or histone modifications (e.g., methylation, acetylation), the latter corresponding to the chromatin state of the region. However, the population genetic effects of these changes are often ignored. Changes to the DNA methylation and chromatin state may affect both the regulatory environment of the genes present (also potentially altering dominance patterns) and the recombination rate. In general, an increase in DNA methylation or heterochromatinization will often decrease recombination (crossovers are less likely in highly heterochromatic regions [Henderson, 2012]) and gene expression.

DNA methylation levels and chromatin state are two factors widely responsible for the regulation of genes, either up- or downregulating certain regions of the genome (Talbert et al., 2019). Among different mutation types, transposable elements (TEs) in particular lead to the alteration of the methylation and chromatin context around themselves. This is because TEs are in a constant arms race with the host, and genomes have evolved multitudes of sequence-specific mechanisms to silence new TE insertions via DNA methylation (e.g., CpG dinucleotides) and repressive histone marks (e.g., H3K9me2 and H3K9me3; Choi & Lee, 2020; Hollister & Gaut, 2009). These changes in methylation and chromatin state not only affect the newly inserted TE itself (and its possible fitness effects) but may spread into adjacent genomic regions, for example, up to 20 kb away from TE insertions in *Drosophila melanogaster* (Lee & Karpen, 2017) acting as a local DFE modifier. Translocations may also change DNA methylation state; a study on humans found multiple differentially methylated positions with respect to a translocation, 93% of which mapped to the translocation breakpoints (McCartney et al., 2018). Finally, chromosomal fusions have been reported to be associated with larger regions bearing repressive histone marks in mice, potentially leading to a decrease in recombination events (Capilla et al., 2014).

Centromere shifts either happen through a change in chromatin state alone ("neocentromeres"; Marshall et al., 2008) or together with the expansion of specific repetitive sequences ("evolutionary new centromeres"; Rocchi et al., 2012). Centromeres contain both centromeric chromatin (characterized by the CENP-A histone, which is the foundation for the kinetochore), and repressive histone marks (Sullivan & Karpen, 2004). Nevertheless, their effects on population genetics parameters can be expected to be similar to other structural variants. The shift of the centromere along a chromosome will directly reduce the recombination rate in the new centromere-adjacent region (see Section 2.1). A reduction in recombination will increase the rate of accumulation of TEs, spreading DNA methylation and repressive chromatin marks indirectly. This generates a positive feedback loop between the reduction of the local recombination rate, new TE insertions, and change in chromatin state as previously proposed for regions of low recombination in general (Kent et al., 2017).

More work remains to be done to determine the effect of large structural variants and other mutations on DNA methylation levels and chromatin state. While DNA methylation levels and chromatin state tend to be less permanent than sequence changes, their consequences are far reaching if they also alter the state of the flanking regions. Furthermore, DNA methylation and chromatin state may play a role in phenotypic adaptation, as such changes can shift phenotypes over short time scales (Carneiro & Lyko, 2020; Pikaard & Mittelsten Scheid, 2014; Stajic et al., 2021). Understanding these effects and their phenotypic consequences will help towards building a more unified framework for analysing the relative role of the various mutation types in evolutionary processes.

BOX 2 From mutation type to population genetic effect: Using theory to quantify the impact of structural variants on recombination

Here, we present an example of the forward approach. We (i) summarize mechanistic information about recombination reduction in some types of large structural variants, and (ii) show how this information can be incorporated theoretically.

Mechanisms of recombination reduction

Recombination in eukaryotes begins with double strand breaks (DSBs) that form during the pairing of the homologues in meiosis and are repaired via two pathways. (i) A crossover event (CO), the outcome of which is visible as a chiasma later in meiosis, or (ii) the break is repaired as a noncrossover (NCO) event. Gene conversion (GC) can occur in both pathways (Hunter, 2015). Direct changes in recombination rate can be due to changes in the pairing process, distribution of recombination events, or pathway taken.

Several mutations affect the alignment and pairing of homologues at the beginning of the recombination process when heterozygous. In inversion heterozygotes, proper synapsis in the inverted region and subsequent crossing over are slightly reduced (Gong et al., 2005). A large heterozygous indel will generally form "unpaired DNA loops" preventing COs (Poorman et al., 1981). Similarly, copy number variants (CNVs) can affect recombination in heterozygotes due to differences in chromosome length, effectively reducing recombination by inhibiting proper pairing (Sjödén & Jakobsson, 2012).

Mutations may also affect the pathway taken after a DSB (i.e., a CO or an NCO). The presence of fusions changes the rates and distribution of chiasmata (indicative of a CO) in both homo- and heterozygotes in a range of mammals (Dobigny et al., 2017). For example, in mice (*Mus musculus domesticus*), the number of chiasmata correlates negatively with the number of fusions but the distribution of the chiasmata along the chromosomal arm varies between homozygotes and heterozygotes (Bidau et al., 2001; Capilla et al., 2014). In inverted regions, DSBs are more likely to be resolved as NCOs; however, the rate of GC is unchanged (Crown et al., 2018; Korunes & Noor, 2019). Conversely, heterozygous translocations do not affect the ratio of COs to NCOs but reduce the rate of GC (Sherizen et al., 2005).

Recombination may occur normally but lead to the creation of unbalanced gametes in heterozygotes only. In inversions, crossing over in the inverted region leads to gametes with unbalanced chromosomes with potentially large duplications and deletions (Rieseberg, 2001). However, there are several mechanisms that can reduce the creation of unbalanced gametes. First, DSBs can be repaired as NCOs (see above) or the rate of DSBs can be reduced in the inverted region, both of these will reduce the production of unbalanced gametes (Fuller et al., 2019). Furthermore, when inversions are heterozygous, the inverted region can either pair homosynaptically or heterosynaptically (Torgasheva & Borodin, 2010). COs can only occur in homosynaptically paired regions so increasing the rate of heterosynaptic pairing reduces the production of unbalanced gametes (Torgasheva & Borodin, 2010). Alternatively, recombination can proceed normally and create balanced products but these products may fail to segregate properly. In translocation heterozygotes, the four involved chromosomes form a quadrivalent structure during meiosis. Segregation from this structure can lead to the creation of aneuploid gametes with a rate of 18% to more than 80% (Morel et al., 2004; Talukdar, 2010). Similarly, nondisjunction rates in fusion heterozygotes may be elevated, ranging from 1.2% to 30% depending on the system (Dobigny et al., 2017).

Theoretical incorporation

To quantify the effects described above, we derive the probability, $P(x_1, x_2)$, that two loci at position x_1 and x_2 (with $x_1 < x_2$; measured in bp), initially on the same homologue, are separated during meiosis in the presence of various structural variants. We present here only approximations obtained when the rate of double strand break (DSB), β_{DSB} , is sufficiently small (see Supporting Information for detailed expressions).

In the absence of a structural variant, the probability that two loci are separated by recombination is given by:

$$P_{\text{rec}}(x_1, x_2) \cong \beta_{\text{DSB}}(\lambda\phi_{\text{GC}} + (x_2 - x_1)\phi_{\text{CO}}),$$

with ϕ_{GC} and ϕ_{CO} as the probabilities that a DSB leads respectively to gene conversion (GC) and a crossover and λ the length of a GC tract ($x_2 > x_1 + \lambda$). The first term corresponds to one locus being transferred by GC and the second to a crossover between the two focal loci.

Insertion/deletion (Indel)

Recombination only happens in the ancestral (deletion) or derived (insertion) homozygote (its frequency denoted f_{AA}). The two loci are separated with probability:

$$P_{\text{Indel}}(x_1, x_2) = f_{AA} P_{\text{rec}}(x_1, x_2).$$

Inversion

Single crossovers occurring within the inversion breakpoints in heterozygotes form gametes with unbalanced chromosomes, leading to inviable zygotes. Here, we depict a case where heterozygotes are underdominant and recombination only happens through GC or double crossovers. The probability that two loci in the inverted region are separated is given by (assuming $\beta_{\text{DSB}} \ll f_{AA} - 0.5$):

$$P_{\text{inv}}(x) \cong \left(\frac{(x_2 - x_1)(1 - 2f_{AA})\phi_{\text{CO}}}{1 - f_{AA}} + \lambda\phi_{\text{GC}} \right) \beta_{\text{DSB}}.$$

The first term corresponds to a recombination event happening between the focal loci in the homozygotes, whose frequency is increased due to underdominant heterozygotes. The second term corresponds to GC and remains unaffected. Double crossovers do not play a significant role under those conditions.

Fusion

For chromosomal fusions, homologues in heterozygotes may fail to segregate properly (with probability β_{NDJ}), producing unbalanced gametes and reducing the contribution of heterozygotes to the next generation. In addition, the chance of a crossover decreases if at least one fused chromosome is involved. The two loci are separated with probability:

$$P_{\text{fus}}(x_1, x_2) \cong (S_1(f_{AA}, f_{AA}))(x_2 - x_1)\phi_{\text{CO}} + \lambda\phi_{\text{GC}}\beta_{\text{DSB}}.$$

The contribution of crossovers is reduced by a factor $S_1(f_{AA}, f_{AA})$, which depends on the genotypes frequencies and captures both selection against the heterozygote and the reduced crossover probability when at least one fused chromosome is involved.

Translocation

Similarly, homologues in heterozygotes may fail to segregate properly, producing unbalanced gametes and reducing the contribution of heterozygotes to the next generation. The GC rate in heterozygotes is also reduced. The two loci are separated with probability:

$$P_{\text{trans}}(x_1, x_2) \cong (x_2 - x_1)\phi_{\text{CO}} + S_2(f_{AA})\lambda\phi_{\text{GC}}\beta_{\text{DSB}}.$$

The contribution of gene conversion is reduced by a factor $S_2(f_{AA})$, which depends on the frequency of the heterozygotes and captures both the effect of selection against, and the reduction of gene conversion within, heterozygotes.

Overall, these results show that the extent of recombination reduction probably differs between mutation types. Although the parameter space remains unexplored, under our assumptions, recombination was most strongly reduced in indels followed by inversions. Surprisingly, recombination in the translocation closely mirrored default recombination rate.

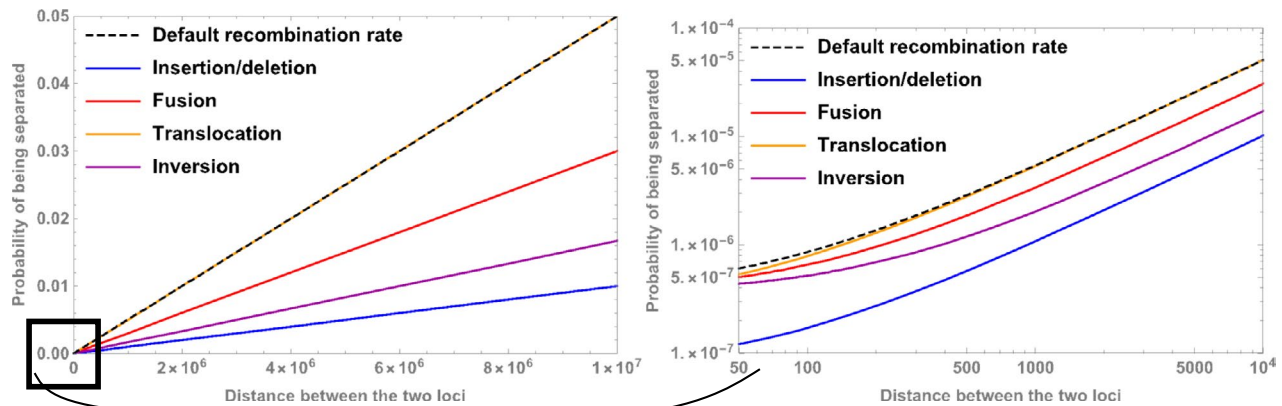


Figure Box 2. Probability that two loci on the same structural variant are separated due to recombination as a function of the distance between the two loci (in bp). Parameters: $\beta_{DSB} = 10^{-8}$; $\phi_{CO} = 0.5$; $\phi_{GC} = 0.7$; $\lambda = 50$; $f_{Aa} = 0.4$; $f_{AA} = 0.2$. The factors S_1 and S_2 are calculated using the expressions given in the Supplement; here $S_1 = -0.6$ and $S_2 = -0.8$

2.1 | Recombination rate and physical distance

The probability that two loci are separated during meiosis is affected by segregation patterns, the physical distance between them (physical linkage), and the per base pair recombination rate. Mutations such as centromere shifts can distort their segregation during female meiosis of heterozygotes (a process known as centromere drive) (Malik, 2009). Other mutations such as fusions and translocations will affect segregation patterns by bringing previously completely (physically) unlinked loci into linkage. Additionally, translocations will also break cosegregation of loci on either side of the breakpoints by moving them to separate chromosomes. Fusions, translocations, inversions and large indels will all affect physical distance between loci (Smukowski & Noor, 2011).

Mutations may also alter the local recombination rate. There is a rich history of population genetic models linking recombination modifiers with evolutionary outcomes (Barton, 1995; Ortiz-Barrientos et al., 2016; Otto & Barton, 1997). For example, a now classic model by Kirkpatrick and Barton (2006) showed that the reduced recombination provided by inversions can be directly selected for and facilitate local adaptation with gene flow. However, the mechanism by which recombination is reduced and the extent of this reduction varies greatly between mutation types and this will have strong downstream consequences for other population genetic effects, such as dominance, which will impact the evolutionary fate of the mutation (see below). Using several types of large structural variants as an example, we explore the relationship between molecular mechanism and recombination rate and their integration in theoretical models in Box 2.

Other variants alter recombination via changes in the chromatin landscape and/or by insertion of specific sequence motifs. Centromere shifts change local recombination by strongly altering

the chromatin environment (Stapley et al., 2017). Centromere shifts along the chromosome reduce recombination in the new centromere-adjacent region while the former inactivated centromere is then free of centromere-associated recombination reduction. To our knowledge, these aspects of centromere evolution have yet to be appreciated in an evolutionary context. Transposable element (TE) insertions are a type of insertions that can actively change recombination rates via chromatin environment or sequence motifs. Their effect depends on whether they attract repressive histone marks locally decreasing recombination (see Box 1; Choi & Lee, 2020) or contain sequence motifs that turn the region into a recombination hotspot (Kent et al., 2017). Similarly, simple sequence repeats (SSRs) can also act as recombination hotspots or recombination repressors (Brandström et al., 2008; Guo et al., 2009; Myers et al., 2005).

2.2 | Dominance

Dominance determines the penetrance of a mutation and its visibility to selection and can have complex effects on evolutionary processes. Previous population genetic models have shown that the fixation probability of a dominant beneficial mutant is higher than when it is recessive (Haldane's sieve) (Haldane, 1927), while the opposite trend holds if the mutant is deleterious (Kimura, 1957). While best examined on a case by case basis, there are a few trends that have been noted considering mutation types and dominance. For example, as long as an insertion contains a single-copy gene (i.e., a gene that is not present elsewhere in the genome), alleles that are normally recessive will be expressed in the heterozygous state. CNVs themselves will alter the penetrance of a dominant mutation. For example, duplications of the recessive allele may nullify the dominant

mutation or compensate identical alleles with low gene expression level (Beckmann et al., 2007). Other mutations may also have effects that spread outside of the mutated region. For example, many TEs contain regulatory elements for their own mobility, potentially rewiring the regulation of nearby genes, altering dominance patterns (Chuong et al., 2017).

The dominance effects of inversions depend on multiple factors. For example, an inversion might be underdominant if COs in the inversion region lead to unbalanced gametes (see above). On the other hand, recessive deleterious alleles can accumulate within both the standard and the inverted arrangement, generating (associative) overdominance at the level of the inversion because recessive alleles are shielded in inversion heterokaryotypes (Ohta, 1971).

2.3 | Effective population size

Effective population size (N_e) usually reflects the process of drift. As a rule of thumb, a mutation is selectively neutral when $|s| < 1/(2N_e)$

thus all forms of selection (directional, purifying, balancing, etc.) are less efficient as N_e decreases. N_e can vary along the genome and can be locally or globally affected by mutations, either by a direct reduction in the number of gene copies or indirectly through the mutation's effects on recombination rate and selection coefficient (Charlesworth, 2009; Gossman et al., 2011).

Recombination and therefore N_e can be reduced by a variety of mutation types. For example, loci within a large indel will experience this reduction in N_e twice, once due to the reduction in recombination (see Box 2) and once due to lower number of copies of the indel region. The indel as a locus, with two alleles ("present" and "absent"), will not be affected by either of these processes. Similarly, recombination between different arrangements of a heterozygous inversion is lowered and the arrangements can be viewed as two smaller and partially isolated populations (Berdan et al., 2021; Faria et al., 2019). Translocations and fusions will experience a similar effect.

Changes in fitness due to mutations can also lead to a reduction in N_e . For example, TE insertions are weakly deleterious in many sequence contexts (Choi & Lee, 2020; Hollister & Gaut, 2009) leading

BOX 3 Contribution of different mutation types to speciation

Here, we use speciation as an example to demonstrate the application of our framework.

(a) *Predicting the contribution of different mutation types.* Recombination is a key population genetic effect relevant for speciation with gene flow. In the presence of gene flow, speciation can only progress when associations (linkage disequilibria) between alleles at different loci contributing to population differences are maintained and increase (Smadja & Butlin, 2011), and theoretical work predicts that such associations are facilitated by structural variants that reduce recombination (Kirkpatrick & Barton, 2006). This effect has been mostly studied for inversions, but other recombination-reducing mutations, including fusions and maybe centromere shifts, might also be important.

As discussed in the main text, the contribution of a mutation type to an evolutionary process is determined not only by its population genetic effects but also by its mutation rate. SNPs are the most commonly occurring types of variants (Figure 1b). While DFE studies have shown that the majority of non-synonymous SNPs are typically deleterious (Eyre-Walker & Keightley, 2007), SNPs are still likely to make a major contribution to divergence and speciation. TEs have also been hypothesised to be particularly relevant here, as their mutation rates can increase under stress. Increased TE activity in new environments might generate novel diversity, some of which may be adaptive and contribute to population divergence (Stapley et al., 2015).

While this Box demonstrates that different mutation types have been predicted to play a role in speciation, their relative importance is less clear. For example, making more detailed predictions about the relative importance of different recombination-reducing mutations requires more empirical data on their effects on recombination, the selection pressures acting on them (e.g., over- vs. underdominance), as well as theoretical models and simulations that directly compare them (Box 2).

(b) *Empirical data about the relative contribution of different mutation types to speciation with gene flow.* There are several well-studied systems where multiple mutation types have been analysed. For example, for stickleback freshwater-marine divergence, causal mutations are known to include SNPs/small indels (Archambeault et al., 2020) as well as deletions (Chan et al., 2010). Furthermore, differentially adapted populations also differ in the frequency of an inversion (Jones et al., 2012). Flowering plants of the *Mimulus* species complex are another example where multiple mutation types, including SNPs, inversions, translocations and duplications, have been studied (Twyford & Friedman, 2015; Zuellig & Sweigart, 2018).

However, studies looking systematically for the relative contribution of all different mutation types are essentially lacking. Part of the problem is that it is often difficult to pinpoint the exact causal mutation, rather than identifying just a larger genomic region associated with population divergence or underlying divergent traits. As these genomic regions typically contain many small variants, it is difficult to determine whether the causal variant is a SNP, indel, or TE, for example. Additional studies looking at repeated evolution and/or functional studies (i.e., CRISPR/Cas9 modification of a single locus) will be necessary to clarify this. Overall, further work is needed to systematically compare the contribution of different mutation types in the same study system.

them to be removed by selection along with linked neutral variation (i.e., background selection). Translocations or chromosome fusions lead to high rates of nondisjunction and subsequent negative selection against heterozygotes reducing their contribution to future generations (Dobigny et al., 2017; Morel et al., 2004; Talukdar, 2010). Conversely, centromere shifts may be under positive selection if they exhibit centromere drive (Malik, 2009).

2.4 | Selection coefficient

All of the population genetic effects described above combined with the functional consequences of the mutation will together regulate the interaction of the mutation with selection and drift, and determine evolutionary outcomes (Box 3). Changes in the interaction with selection and drift are partially quantified in the selection coefficient, a measure of differences in relative fitness, encompassing multiple population genetic effects. The selection coefficient of a mutation depends on a multitude of genomic factors including (i) the genomic context, for example, whether it alters coding, regulatory, or neutrally evolving regions; and (ii) whether or not it causes a positional shift; but also on nongenomic factor such as the selective environment (both extrinsic and intrinsic) where the change occurs (Brandström et al., 2008; Crown et al., 2018; Ducos et al., 2007; Flynn et al., 2020; Gemayel et al., 2010; Guo et al., 2009; Hollister & Gaut, 2009; Kayser et al., 2006; Kent et al., 2017; Korunes & Noor, 2019; Sherizen et al., 2005; Stapley et al., 2017; Weissensteiner et al., 2020). Furthermore, duplicated regions, such as CNVs, have additional effects as they may free up selective constraints and can lead to the emergence of new gene functions (Ohno, 2013).

Selection coefficients can be examined more globally using the distribution of fitness effects (DFE), that summarizes the interaction of the mutation type with drift and selection (Eyre-Walker & Keightley, 2007; Keightley & Eyre-Walker, 2010). There is a large body of work describing the statistical properties of the DFE both during adaptation and for well adapted populations, mostly within the framework of Fisher's Geometric Model (FGM: Allen Orr, 2005; Tenaillon, 2014). FGM predicts that the size of the effect of fixed mutations decreases as the populations become adapted to their environment. This prediction has been thoroughly tested and there are empirical studies that show results compatible with the FGM (Blanquart & Bataillon, 2016; Harmand et al., 2017; Martin et al., 2007; Martin & Lenormand, 2006). However, the mechanistic underpinnings of this change in effect size across adaptation is unknown. A potential explanation is the fixation of different types of mutations at different stages of the adaptive process, something, that to the best of our knowledge, has not been yet tested, either empirically or theoretically. However, since FGM is a phenotype-based model, it makes no assumptions about the type of mutations underlying the different mutational steps. Linking the FGM with genomic data is a promising avenue to better understand the DFE of different mutation types and connect this with adaptive processes.

Most studies have estimated the DFE of SNPs and have found a bi- or multimodal distribution, with beneficial mutations being rare, although the exact shape of distributions vary (Bataillon & Bailey, 2014; Eyre-Walker & Keightley, 2007; Keightley & Eyre-Walker, 2010). However, the DFE of other mutation types may have different properties (but see Barton & Zeng, 2018). For example, a study in *E. coli* (Elena et al., 1998) showed a long deleterious tail and a high neutral peak for TE insertions. Most CNVs that have been characterized were found at the extremes of the distribution with either beneficial or largely deleterious effects (Katju & Bergthorsson, 2013). However, these studies are biased towards large effect CNVs so the true distribution remains unknown. While the DFE allows us to make certain evolutionary predictions it does not quantify downstream effects of critical population genetic effects such as recombination rate or genomic effects such as changes in the chromatin landscape that may alter the evolutionary trajectory of the mutation.

Large structural variants alter the efficacy of selection within the mutated region by modifying the recombination rate and local N_e . This aspect can be beneficial; for example, inversions may be indirectly selected because they reduce recombination between multiple beneficial alleles located in the same arrangement (e.g., locally adapted alleles under gene flow [Kirkpatrick & Barton, 2006]). However, these changes also alter the evolution of the mutated region in multiple ways, for example, by reducing the efficacy of purifying selection leading to the accumulation of deleterious alleles. Quantifying the impact of these changes, through a combination of analytical approaches and simulations (e.g., Gilbert et al., 2020) will be a key step towards linking mutation type with evolutionary significance.

3 | POPULATION GENETIC EFFECTS OF MUTATIONS: CONNECTING TO EVOLUTIONARY OUTCOMES

To understand the evolutionary significance of a mutation, we must examine its past, current and potential future effects. Our framework outlines a combination of theoretical and empirical tools that can be used to: (i) Estimate the population genetic effects of all mutation types simultaneously, (ii) determine how these effects can shape the evolutionary trajectory of the population, and (iii) combine 1 and 2 to link these effects to evolutionary outcomes (e.g., speciation, adaptation).

As summarized above, there is already a large body of work describing the molecular underpinnings of a mutation's population genetic effects (Katju & Bergthorsson, 2013; Kirkpatrick, 2010). However, while we sometimes know the general direction of these effects (Figure 1c), we do not know much about their effect size distributions. Empirical studies quantifying the population genetic effects of mutations on a large scale are sorely needed for better characterization of mutation effects and to determine how these effects vary across taxa. The DFE is a good place to start, but we also need to quantify the other population genetic effects of mutations.

A critical question is how these effect sizes vary between mutation types. For example, to what extent is recombination reduced in an inversion compared to a fusion? This information can be directly related to evolutionary outcomes, e.g., is a fusion or an inversion a “better” genomic architecture for maintaining a complex of coadapted alleles? With enough data it may even be possible to estimate distributions of population genetic effects for different mutation types. Below we discuss both empirical and theoretical ways forward.

Several of the population genetic effects described here are relatively straightforward to measure in model systems. For example, the influence of a mutation on recombination can be determined using mapping crosses followed by sequencing and bioinformatic detection of gene conversion and crossing over events. The effect of inversions on recombination in *Drosophila* has already been intensively examined using this methodology (Crown et al., 2018; Korunes & Noor, 2019). The application of these methods to other systems and other types of mutations (e.g., fusions) will allow us to determine the distribution of recombination suppression for different types of mutations. Changes in physical distance can be quantified in part by utilizing pre-existing sequence data to examine the size distribution for different types of structural mutations. These types of studies provide a start for building distributions of population genetic effects.

In addition to the empirical approaches described in the last paragraph, we can begin to ask this question using theoretical models. We provide an example in Box 2 showing how molecular genetic information about recombination may be incorporated theoretically. Although exploring the parameter space was beyond our scope, these results illustrate that the extent of recombination reduction probably differs between mutation types. Box 2 represents a starting point for integrating underlying molecular mechanisms with their impacts on population genetic parameters. Complementing analytical approaches with simulations allows for exploration of more complex effects on a larger scale.

In order to determine how these population genetic effects link to evolutionary outcomes we must examine how they shift evolutionary trajectories. Theoretical models provide the best avenue for this. There is more than a century of literature developing these methods in population genetics (Box 3, Charlesworth & Charlesworth, 2010). Integrating the feedback loop between the evolution of a structural variant as a locus and the evolution of its allelic content into theoretical models may further our understanding of the link between mutation types and evolutionary outcomes. For example, in the case of an inversion, the resulting reduction in recombination rate generally leads to an accumulation of deleterious alleles in the minor variant, slowly increasing the fitness differential with the major common variant (Berdan et al., 2021). Looking at empirical data can show the result of these shifts in trajectory. Patterns of nucleotide diversity, divergence between arrangements, and the DFE of the alleles within the mutated region can be examined and compared with predictions from simulation studies. In this way, the forward and reverse approaches can be merged (Figure 1a).

Empirical studies also provide critical information about what mutation types have previously been important in evolutionary outcomes. However, most studies do not compare different types of mutations. Moving forward will require collecting different types of genomic data sets (e.g., short- and long-read re-sequencing and mapping crosses) from the same population and developing detection pipelines targeted at different mutation types (Mérot et al., 2020). Additionally, new computational methods expedite the identification of different types of mutations, for example using machine learning (Caravagna et al., 2020) and parallel computation (Larson et al., 2019), combining multiple statistical models (Huang et al., 2017) and integrating different types of sequencing (Harmanci et al., 2020; Sobreira et al., 2011). Synthesizing information on mutation types and evolutionary outcomes allows us to both explore the relationships between mutation type and the major evolutionary outcomes and to test predictions based on population genetic effects. For example, speciation requires the build-up of linkage disequilibrium between alleles contributing to reproductive isolation (Box 3, Butlin & Smadja, 2018). Mutations that reduce recombination should aid speciation with gene flow by protecting this nascent linkage disequilibrium. We can thus predict that mutations such as inversions, large indels, TEs, fusions, and centromere shifts might be major drivers of speciation events (Fuller et al., 2018). A critical next step would be testing some of these hypotheses in a quantitative rather than review framework, for example using a meta-analysis.

Experimental evolution offers another way to integrate the forward and reverse approaches detailed above. These studies link mutation type with evolutionary outcome in real time (Kawecki et al., 2012), generating results that can be compared with theoretical predictions and empirical results from natural populations. For example, starting populations for experimental evolution studies can incorporate genetic variation for multiple mutation types (e.g., segregating inversions and CNVs). The evolutionary trajectories of these different mutation types can then be followed during the adaptive or divergence process and these can be combined with functional studies to pinpoint adaptive variants. In this way the relative role of different mutations can begin to be dissected. Concomitantly, existing genomic data from previous experimental evolution studies can also be utilized. By using different software programs (Chen et al., 2016; Kawecki et al., 2012; Liu et al., 2020; Moreno-Cabrera et al., 2020; Shigemizu et al., 2018) to detect different types of mutations, it should be possible to infer the relative role of different mutations in these different scenarios. Overall, experimental evolution studies can provide a valuable counterpoint to theoretical predictions and data from more traditional population genomic studies (Moreno-Cabrera et al., 2020).

4 | CONCLUDING REMARKS AND FUTURE PERSPECTIVES

Our framework highlights the fact that each mutation type may affect evolution in several ways and that many different mutation types have similar population genetic effects (Box 2, Figure 1).

Analysing this in a quantitative and comparative way will allow us to explore the evolutionary significance of different mutation types.

Understanding the relative evolutionary significance of different mutations will require viewing their effects in a larger population genetic context. In order to do this we need: (i) Comparable measurements of mutation rates as well as the population genetic effects of different mutation types; (ii) to include these effects in theoretical models and simulations to create predictions about the importance of different mutation types; and (iii) to empirically estimate the contributions of different mutations to evolutionary outcomes and test the predictions obtained from the theoretical models. Superimposing a more integrated framework on previous and future work will allow us to better understand the relative contributions of different mutation types to key evolutionary outcomes, further illuminating the genetic underpinnings of these processes in a broad sense.

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AUTHOR CONTRIBUTIONS

Inês Fragata and Emma L. Berdan conceived the study. Inês Fragata, Alexandre Blanckaert, Alexander Suh, Tanja Slotte, Emma L. Berdan and Anja M. Westram wrote the manuscript.

DATA AVAILABILITY STATEMENT

This is an opinion article and does not contain data.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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